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- (54) Homogeneous olefin polymerization catalyst by ligand abstraction with lewis acids.
- Addition polymerization catalysts having a limiting charge separated structure corresponding to the formula:
 LMX+ XA-,

wherein :

L is a derivative of a substituted delocalized Π -bonded group imparting a constrained geometry to the metal active site and containing up to 50 nonhydrogen atoms;

M is a metal of Group 4 or the Lanthanide series of the Periodic Table of the Elements;

X independently each occurrence is hydride or a hydrocarbyl, silyl or germyl group having up to 20 carbon, silicon or germanium atoms; and

A⁻ is an anion of a Lewis acid, A, having relative Lewis acidity greater than or equal to that of phenylbis(perfluorophenyl)borane, said anion being compatible with the metal cation, are prepared by contacting a derivative of a Group 4 or Lanthanide metal corresponding to the formula:

LMX₂,

wherein

L, M, and X are as previously defined, with the Lewis acid, A.

This invention relates to a method for preparing compositions of matter that are useful as addition polymerization catalysts. More particularly, this invention relates to a method for preparing homogenous, constrained geometry, cationic, catalyst compositions by abstraction with Lewis acids.

In EP-A 416,815 ther are disclosed and claimed certain constrained geometry metal complexes having utility as homogeneous olefin polymerization catalysts. In EP-A-418,044 cationic monocyclopentadienyl m tal complexes with salts of Bronsted acids containing a non-coordinating compatible anion are disclosed and claimed. In EP-A-468,651, an oxidative activation technique for preparing such cationic catalysts is disclosed and claimed.

In <u>J. Am. Chem. Soc.</u> 113, 3623-3625 (1991) there is disclosed a process for preparation of "cation like" zirconocene polymerization complexes by alkyl abstraction using tris(pentafluorophenyl)borane. The complexes were stated to have activity roughly comparable to typical complexes based on alumoxane. No suggestion as to the suitability of the disclosed technique for use with constrained geometry metal catalysts is provided by the reference.

In J. Organometal. Chem., 22, 659-663 (1970), there is disclosed the reaction between tetramethyl-titanium and triphenylborane or tribenzylborane. The authors postulate the intermediate formation of cationic titanium species which were not isolated during the reaction. No suggestion as to the suitability of the disclosed reaction for use in catalysis is provided by the reference.

Previously known techniques for preparing cationic metal complexes having constrained geometry have been disadvantaged by requiring difficult synthetic procedures to prepare the necessary precursors and low yields of isolated products. It would be desirable if there were provided an improved technique for preparing cationic metal complexes having constrained geometry which avoids difficult synthetic procedures and poor vields.

It has now been discovered that the foregoing and other disadvantages of the prior art cationic, olefin polymerization catalysts can be avoided or at least reduced with the process of the present invention. Surprisingly the catalysts of the present invention are marked by extremely high catalytic effectiveness as measured by polymer yield at a given temperature.

In accordance with the present invention there is provided a process for preparing a catalyst having a limiting charge separated structure corresponding to the formula:

LMX+ XA-,

30 wherein:

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L is a derivative of a substituted delocalized Π -bonded group imparting a constrained geometry to the metal active site and containing up to 50 nonhydrogen atoms;

M is a metal of Group 4 or the Lanthanide series of the Periodic Table of the Elements;

X independently each occurrence is hydride or a hydrocarbyl, silyl or germyl group having up to 20 carbon, silicon or germanium atoms; and

A⁻ is an anion of a Lewis acid, A, having relative Lewis acidity greater than or equal to that of phenyl-bis(perfluorophenyl)borane, said anion being compatible with the metal cation,

the steps of the process comprising contacting a derivative of a Group 4 or Lanthanide metal corresponding to the formula:

LMX₂,

wherein

L, M, and X are as previously defined,

with the Lewis acid, A.

The preceding formula for the catalyst compositions is referred to as the limiting, charge separated structure. However, it is to be understood that, particularly in solid form, the catalyst may not be fully charge separated. That is, the X group may retain a partial covalent bond to the metal atom, M. Thus, the catalysts may be alternately depicted as an ion pair possessing the formula:

LMX--X--A.

The catalysts are preferably prepared by contacting the derivative of a Group 4 or Lanthanide metal with the Lewis Acid in an inert diluent such as an organic liquid.

All reference to the Periodic Table of the Elements herein refer to the Periodic Table of the Elements, published and copyrighted by CRC Press, Inc., 1989. Also, any reference to a Group or Groups are to the Group or Groups as reflected in this Periodic Table of the Elements using the IUPAC system for numbering groups.

By use of the term "constrained geometry" herein is meant that the metal atom is forced to greater exposure of the active metal site because n or mor substituents on th substituted delocalized Π -bonded group forms a portion of a ring structure including the m tal atom, wherein th m tal is both bonded to an adjacent covalent moiety and held in association with the substituted delocalized Π -bonded group through an η^5 or other Π -bonding interaction. It is understood that each respective bond between the metal atom and th constituint atoms

of the substitut d delocalized Π -bonded group need not b equivalent. That is, the metal may be symmetrically or unsymmetrically Π -bonded to the substituted delocalized Π -bonded group.

The geometry of the active metal site is further defined as follows. The center of the substituted delocalized Π-bonded group may be defined as the average of the respective X, Y, and Z coordinates of the atomic centers forming the substituted delocalized Π-bonded group. The angle, if med at the metal center between the center of the ligating atom of each other ligand of the metal complex may be easily calculated by standard techniques of single crystal X-ray diffraction. Each of these angles may increase or decrease depending on the molecular structure of the constrained geometry metal complex. Those complexes wherein one or more of the angles, Θ, is less than in a similar, comparative complex differing only in the fact that the constrain-inducing substituent is replaced by hydrogen have constrained geometry for purposes of the present invention. Preferably one or more of the above angles, Θ, decrease by at least 5 percent, more preferably 7.5 percent, compared to the comparative complex. Highly preferably, the average value of all bond angles, Θ, is also less than in the comparative complex.

Preferably, metal coordination complexes of Group 4 or Lanthanide metals according to the present invention have constrained geometry such that the smallest angle, Θ , is less than 115°, more preferably less than 110°, most preferably less than 105°.

Substituted delocalized Π -bonded groups for use herein include any Π -electron containing moiety capable of forming a delocalized bond with the Group 4 or Lanthanide metal and further substituted with one or more divalent substituents that are also covalently bound to the metal. Divalent substituents preferably include groups containing up to 30 nonhydrogen atoms comprising at least one atom that is oxygen, sulfur, boron or a member of Group 14 of the Periodic Table of the Elements directly attached to the delocalized Π -bonded group, and a different atom, selected from nitrogen, phosphorus, oxygen and sulfur that is covalently bonded to M. Examples of suitable delocalized, Π -bonded groups are cyclopentadienyl- or allyl-groups, and derivatives thereof.

By the term "derivative" in the definition of L is meant that each atom in the delocalized Π -bonded group may independently be substituted with a radical selected from hydrocarbyl radicals, substituted-hydrocarbyl radicals wherein one or more hydrogen atoms are replaced by a halogen atom, hydrocarbyl-substituted metalloid radicals wherein the metalloid is selected from Group 14 of the Periodic Table of the Elements, and halogen radicals. Suitable hydrocarbyl and substituted-hydrocarbyl radicals used to form derivatives of the delocalized Π -bonded group contain from 1 to 20 carbon atoms and include straight and branched alkyl radicals, cyclic hydrocarbon radicals, alkyl-substituted aromatic radicals. In addition two or more such radicals may together form a fused ring system or a hydrogenated fused ring system. Examples of the latter are indenyl-, tetrahydroindenyl-, fluorenyl-, and octahydrofluorenyl- groups. Sultable hydrocarbyl-substituted organometalloid radicals include mono-, di- and trisubstituted organometalloid radicals of Group 14 elements wherein each of the hydrocarbyl groups contains from 1 to 20 carbon atoms. More particularly, suitable hydrocarbyl-substituted organometalloid radicals include trimethylsilyl, triethylsilyl, ethyldimethylsilyl, methyldiethylsilyl, triphenylgermyl, and trimethylgermyl.

Preferably M is a metal of Group 4 of the Periodic Table of the Elements, most preferably titanium or zirconium. Also, preferably X is C_1 - C_{10} hydrocarbyl, most preferably benzyl and especially methyl.

Highly preferred Group 4 or Lanthanide metal derivatives are substituted monocyclopentadienyl compounds corresponding to the formula:

wherein:

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M is titanium or zirconium:

Cp is a cyclopentadienyl group or derivative thereof that is Π-bound to M and substituted at least by Z; Z is a divalent moiety comprising oxygen, sulfur, boron, or a member of Group 14 of the Periodic Table of the Elements;

Y is a ligand group comprising nitrogen, phosphorus, oxygen or sulfur or optionally Z and Y together form a fused ring system; and

X is as previously defined.

After abstraction of thi X group, the highly preferred catalysts of the invintion have a limiting, electron separated structur corresponding tith formula:

wherein Cp. Z, M, X and A are as previously defined.

In the most preferred embodiment -Z-Y- is an amidosilane or amidoalkane group, preferably of up to 20, especially up to 10, nonhydrogen atoms and especially (tert-butylamido) (dimethylsilyl) or (tert-butylamido)-1-ethane-2-yl.

Most highly preferred derivatives of Group 4 or Lanthanide metals are amidosilanediyl- or amidoalkanediyl-compounds corresponding to the formula:

wherein:

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M is titanium or zirconium, bound to an η⁵-cyclopentadienyl group;

R' each occurrence is independently selected from hydrogen, silyl, alkyl, aryl and combinations thereof having up to 20 carbon or silicon atoms, and optionally two or more R' groups on the cyclopentadienyl group form a fused ring system;

E is silicon or carbon;

X independently each occurrence is hydride, alkyl, aryl or halogen substituted aryl of up to 20 carbons; and

m is 1 or 2.

Examples of the above most highly preferred metal coordination compounds include compounds wherein the R' on the amido group is methyl, ethyl, propyl, butyl, pentyl, hexyl, (including branched and cyclic isomers), norbornyl, benzyl, or phenyl; the cyclopentadienyl group is cyclopentadienyl, indenyl, tetrahydroindenyl, fluorenyl, tetrahydrofluorenyl, or octahydrofluorenyl; R' on the foregoing cyclopentadienyl groups each occurrence is hydrogen, methyl, ethyl, propyl, butyl, pentyl, hexyl, (including branched and cyclic isomers), norbornyl, benzyl, or phenyl; and X is methyl, neopentyl, trimethylsilyl, norbornyl, benzyl, methylbenzyl, phenyl, or pentafluorophenyl.

Illustrative derivatives of Group 4 or Lanthanide metals that may be employed in the practice of the present invention include: [dimethyl(tert-butylamido)(tetramethyl-n5-cyclopentadienyl)silane] dibenzylzirconium, [dimethyl(tert-butylamido)(tetramethyl-n6-cyclopentadienyl)silane] dibenzyltitanium, [dimethyl(tert-butylamido) (tetramethyl-η5-cyclopentadienyl)silane] dimethyltitanium, [(tert-butylamido)(tetramethyl-η5-cyclopentadienyi)-1,2-ethanediyi] dimethylzirconium, [(tert-butylamido)(tetramethyl-η⁵-cyclopentadienyi)-1,2-ethanediyi] dibenzyltitanium, [(methylamido)(η⁵-cyclopentadienyl)-1,2-ethanediyl] dibenzhydrylzirconium, [(methylamido)(tetramethyl-ηδ-cyclopentadienyl)-1,2-ethanediyl] dineopentyltitanium, [(phenylphosphido)(tetramethyl-ηδcyclopentadienyl)methylene] diphenyltitanium, [dibenzyl(tert-butylamido)(tetramethyl -n5-cyclopentadienyl)silane] dibenzylzirconium, [dimethyl(benzylamido)(n5-cyclopentadienyl)silane] di(trimethylsilyl)titanium, [dimethyl(phenylphosphido)(tetramethyl-n5-cyclopentadienyl)silane] dibenzylzirconium, [dimethyl(tert-butylamido)(tetramethyl-n⁵-cyclopentadienyl)-silane) dibenzylhafnium, [(tetramethyl-n⁵-cyclopentadienyl)-1,2-ethanediyl] dibenzyltitanium, [2-ŋ5-(tetramethylcyclopentadienyl)-1-methyl-ethanolato (2-)] dibenzyltitanium, [2-ŋ5-(tetramethylcyclopentadienyl)-1-methyl-ethanolato (2-)] dimethyltitanium, $[2-\eta^5]$ (tetramethylcyclopentadienyl)-1-methyl-ethanolato (2-)] dibenzylzirconium, [2-η5-(tetramethylcyclopentadienyl)-1-methyl-ethanolato (2-)] dimethylzirconium, [2-[(4a, 4b, 8a, 9, 9a -η)-9H-fluoren-9-yl]cyclohexanolato (2-)] dimethyl titanium, [2-[(4a. 4 b, 8a, 9, 9a -η)-9H-fluoren-9-yl]cycloh xanolato (2-)] dibenzyl titanium, [2-[(4a, 4b, 8a, 9, 9a -η)-9H-flu ren-9yi]cyclohexanolato (2-)] dimethyl zirconium, and [2-[(4a, 4b, 8a, 9, 9a -η)-9H-fluoren-9-yi]cyclohexanolato (2-)] dibenzyl zirconium.

Other compounds which ar useful in the catalyst compositions of this invention, especially compounds

containing other Group 4 or Lanthanide metals, will, of course, be apparent to those skilled in the art.

Suitable anions are those that either do not coordinate to or are only weakly coordinated to said metal cation, thereby remaining sufficiently labil t b displaced by a neutral Lewis base. A "compatible" anion additionally ref rs t such an ani n that, when functioning as a charge balancing anion in the catalyst system of this invention, within the time frame of the desired end use, does not transfer an anionic substituent or fragment thereof to the cation thereby forming a neutral metallocene and a neutral metal byproduct. In addition such anions are not degraded to neutrality when the initially formed complex decomposes and are noninterfering with desired subsequent polymerizations or other uses of the complex.

The Lewis acids having Lewis acidity suitable for use according to the present invention are prepared according to known techniques such as those of Marks, et al. <u>J. Am. Chem. Soc.</u> 1991, 113, 3623-3625, or J.Pohlman, et al., <u>Naturforschg.</u> 1965, 20b, 5-11. The preferred technique is to combine a boron or aluminum halide compound such as boron trichloride or boron trifluoride with an alkali metal or alkaline earth derivative of the desired substituent or substituents. Additionally, borate esters such as tris(perfluorophenyl) borate may be prepared by the reaction of pentafluorophenylphenol and borane-methyl sulfide complex according to the method of J. Org. Chem., 43(13) 2731-32 (1978).

Lewis acidity may be measured by empirical means and may also be predicted with a high degree of certainty based on theoretical methods. A preferred technique for measuring Lewis acidity is the proton NMR chemical shift of the proton on carbon #3 (H-3) of crotonaldehyde complexes of the Lewis acid. This technique was disclosed by R. Childs et al., <u>Can. J. Chem.</u>, 1982, 802-808. A similar technique for determining Lewis acidity was taught in P. Laslo et al. <u>J. Am. Chem. Soc.</u> 1990, 12, 8750-8754. The units of measurement are $\Delta\delta$ (ppm). It has been discovered that the technique may be practiced at temperatures at or less than 25°C without deleterious effect.

The difference in chemical shift of the 3-hydrogen of free, uncomplexed crotonaldehyde and the 3-hydrogen of the complexed Lewis acid adduct is determined. This chemical shift difference ($\Delta\delta$ in ppm) is related to the Lewis acidity of the species under study, with the trend being the more downfield the 3-hydrogen is shifted, the greater the Lewis acidity of the compound being examined. The chemical shift difference of phenylbis(perfluor-ophenyl)borane is 0.77 ppm. More Lewis acidic compounds have chemical shift differences, $\Delta\delta$, more than 0.77. Preferred Lewis acids have an acidity from 0.77 to 1.49, more preferably from 1.0 to 1.49. Thus, by the Childs et al. technique, useful Lewis acids for the present invention are acids having relative acidity compared to phenylbis(perfluorophenyl)borane, $\Delta\delta'$ - $\Delta\delta'$, \geq 0 (where $\Delta\delta'$ is the Lewis acidity of the candidate Lewis acid and $\Delta\delta'$ is the Lewis acidity of phenylbis(perfluorophenyl)borane). Preferred Lewis acids are those having relative acidity from 0 to 0.72, more preferably from 0.23 to 0.72.

Detrimental reactivity of Lewis acids includes abstraction of one or more groups from the anion by the cationic portion of the catalyst, LMX+. Readily extractable groups comprise the halides when such halide is directly attached to a central Group 13 metalloid. Thus, most preferred non-reactive Lewis acids are Lewis acids that are devoid of halide groups directly attached to a central Group 13 metalloid, especially boron. Stated otherwise, most preferred Lewis acids are boron compounds devoid of halogen moleties directly attached to the boron.

Theoretical techniques may also be utilized to calculate the acidity of Lewis acids suitable for use according to the present invention. Several commercially available computer programs may be used to calculate the Lewis acidity. In a preferred technique theoretical structures and total energies may be calculated for candidate molecules in a proposed reaction of a Lewis acid with a Lewis base to form a complex. Molecules giving larger calculated heats of complexation indicate greater Lewis acidity. A program such as GAUSSIAN 90, or similar molecular simulation software may be used for the simulation and analysis of such materials.

First, the initial structures are optimized by minimizing the calculated total energy with respect to all degrees of freedom: bond lengths, bond angles, and torsional angles. The heat of reaction (ΔH) is then calculated as being the difference between the total energy of the products and the total energy of the reactants e.g.,

where

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E is approximated by the quantum mechanical energy (E_{QM}) at absolute zero temperature (0° Kelvin) for reactants and products.

By the foregoing technique the Lewis acidity of a compound towards a Lewis base such as methyl anion (CH₃-) or ammonia can be calculated, using the formula:

$$A + (C_6H_5)(C_6F_5)_2B$$
:base $\rightarrow A$:base $+ (C_6H_5)(C_6F_5)_2B$

where A is the Lewis acid and "base" is the Lewis base. If the reaction is exothermic ($\Delta H < 0$), then A is a stronger Lewis acid than phenylbis(perflu rophenyl)boron. Relative acidity is d termined by comparison to the calculated ΔH f phenylbis(perfluorophenyl)boron, which by definition is 0.0 kcal/mol

Based on the foregoing empirical and theoretical t chniques highly preferred Lewis acids are: tris(pentafluorophenyl)boran , tris (2,3,5,6-tetrafluor phenyl)borane, tris(2,3,4,5-tetraflu rophenyl)borane.

tris(3,4,5-trifluorophenyl)borane, tris(1,2,2-trifluoroethenyl)boran , phenylbis(p rfluoroph nyl)boran , tris(3,4,5-trifluorophenyl)aluminum, tris(perfluorophenyl) borate, 1,3,5-cyclohexanetriol borate (cyclohexane-r-1, c-3, c-5-triol borate), and 1,1,1-trimethylolethane borate (2,6,7-trioxa-1-bora-4-m thylbicycl [2.2.2] octan) (the latter two compounds may b prepared according to the conduction of 1,3,5-cyclohexanetriol or 1,1,1-trimethylolethane with b ric acid).

Without wishing to be bound by any particular theory of operation it is believed that the Lewis acid causes the abstraction of the X group and in the process becomes an anionic species. This result is believed to be of practical significance for the present invention where the Lewis acid is a stronger Lewis acid than the resulting cation, LMX*. The specific Lewis acids of the present invention are highly effective in this regard.

In general, the catalyst can be prepared by combining the two components (the derivative of a Group 4 or Lanthanide metal and the Lewis acid) in a suitable solvent at a temperature within the range from -100°C to 300°C, preferably 25 to 50°C. Suitable solvents include straight and branched-chain hydrocarbons such as isobutane, butane, pentane, hexane, heptane, octane, and mixtures thereof; cyclic and alicyclic hydrocarbons such as cyclohexane, cycloheptane, methylcyclohexane, methylcycloheptane, perfluorinated hydrocarbons such as perfluorinated C₄₋₁₀ alkanes and aromatic and alkyl-substituted aromatic compounds such as benzene, toluene and xylene. Suitable solvents also include liquid olefins which may act as monomers or comonomers including ethylene, propylene, butadiene, cyclopentene, 1-hexene, 3-methyl-1-pentene, 4-methyl-1-pentene, 1,4-hexadiene, 1-octene, 1-decene, styrene, divinylbenzene, allylbenzene and vinyltoluene (including all isomers alone or in admixture). Preferred solvents are allphatic hydrocarbons especially C₅-C₁₀ alkanes or cycloalkanes and mixtures thereof.

The catalyst may be used to polymerize addition polymerizable monomers having from 2 to 1000 carbon atoms. Examples include: ethylenically unsaturated compounds, acetylenic compounds, conjugated or non-conjugated dienes, polyenes, and carbon monoxide. Preferred addition polymerizable monomers are olefins or diolefins having from 2 to 18 carbon atoms. Preferred monomers include the C_{2-18} α -olefins especially ethylene, propylene, isobutylene, 1-butene, 1-hexene, 4-methyl-1-pentene, 1-octene, 1-nonene, 1-decene, 1-decene, 1-tetradecene, 1-hexadecene, and 1-octadecene. Other preferred monomers include styrene, halo-or alkyl substituted styrenes, tetrafluoroethylene, vinylbenzocyclobutane, 1,4-hexadiene, norbornene, and substituted norbornenes such as 5-ethylidene-2-nonbornene. Most preferred monomers are α -olefins having from 2 to 12 carbon atoms either alone or in combination. The catalyst may also be used to polymerize α -olefins, diolefins and/or acetylenically unsaturated monomers in combination with other unsaturated monomers.

In general, the polymerization may be accomplished at conditions well known in the prior art for Ziegler-Natta or Kaminsky-Sinn type polymerization reactions, that is, temperatures from 0 to 250°C and pressures from atmospheric to 1000 atmospheres (100 MPa). Suspension, solution, slurry or other process condition may be employed but preferably the catalysts are used in a homogeneous manner. It will, of course, be appreciated that the catalyst system will form in situ if the components thereof are added directly to the polymerization process and a suitable solvent or diluent, including condensed monomer, is used in said polymerization process. It is, however, preferred to form the catalyst in a separate step in a suitable solvent prior to adding the same to the polymerization mixture.

It is believed that the active catalyst species of the present invention contains a metal center which center remains cationic, unsaturated and has a metal-carbon bond which is reactive with olefins, diolefins and acetylenically unsaturated compounds. Also associated with this metal center is a charge balancing anionic remnant of the formula XAT. As previously mentioned the catalyst may also exist in a less than fully charge-separated configuration.

The catalyst formed by the method of this invention may be retained in solution or separated from the solvent, isolated, and stored for subsequent use. As previously indicated supra, the catalyst may also be prepared in situ during a polymerization reaction by passing the separate components into the polymerization vessel where the components will contact and react to produce the catalyst.

The equivalent ratio of derivative of a Group 4, or Lanthanide metal compound to Lewis acid employed is preferably in a range (complex: Lewis acid) from 0.1:1 to 10:1, more preferably from 0.2:1 to 5:1, most preferably 0.5:1.0 to 1:2. In most polymerization reactions the equivalent ratio of catalyst:polymerizable compound employed is from 10⁻¹²:1 to 10⁻¹:1, more preferably from 10⁻⁸:1 to 10⁻⁴:1.

A beneficial feature of the catalysts of this invention, especially the titanium based catalysts, is that when the catalysts of this invention are used to copolymerize olefins, either alone or in combination with diolefins, the amount of higher molecular weight olefin or diolefin incorporated into the copolymer is significantly increased when compared to copolymers prepared with the more conventional Ziegler-Natta typicatalysts. In addition, when compared to constrained geometry catalysts containing alumixane, the catalysts of this present invention possess higher activities. Typically, catalyst efficiencies of the present catalysts (measured by weight

of polymer produced/catalyst weight) are at least fiv times greater than that btained with comparabl alumoxane containing catalysts.

In g neral, catalysts can be select ds as to produce polymer products which will be relatively fr of large amounts of certain trace impurities such as aluminum, magnesium and chlorid g nerally found in polymers produced with Ziegl r-Natta type catalysts. The polymer products produced with the catalysts f this inv ntion have a broader range of applications than polymers produced with more conventional Ziegler-Natta type catalysts comprising a halogenated metal alkyl such as magnesium chloride, or an alkyl aluminum chloride.

Having described the invention the following examples are provided as further illustration thereof and are not to be construed as limiting. Unless stated to the contrary all parts and percentages are expressed on a weight basis.

General Polymerization Procedure

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Ethylene, propytene, and hydrogen were purified by passing through columns of 13X molecular sleve, activated alumina, and trace oxygen removal catalyst (alumina supported copper/manganese catalyst available from Englhardt Inc. under the tradename Q-5). Solvents and octene were degassed with nitrogen, then were purified by passing through columns of 13X molecular sleve, activated alumina, and trace oxygen removal catalyst. Phenylacetylene-free styrene was degassed with nitrogen, then purified by passing through a column of activated alumina. A 2 L stirred autoclave was charged with the desired amounts of a solvent and comonomer.

Hydrogen was added by differential pressure expansion from a 75 mL addition tank. The hydrogen pressure differential represents the difference in pressure between the starting and final pressure in the hydrogen addition tank after adding hydrogen to the 2L reactor containing solvent and comonomer. The reactor was heated to the polymerization temperature and was saturated with ethylene to the desired pressure. Metal complex and Lewis acid cocatalyst were mixed in a drybox by syringing the desired amount of 0.0050 M metal complex solution (in mixed alkane solvent (Isopar E™ available from Exxon Chemicals Inc.) or toluene) into a solution of the cocatalyst (in Isopar® E or toluene solvent). Optionally, additional solvent was added to facilitate transfer of the catalyst solution to the reactor. This solution was then transferred to a catalyst addition tank and injected into the reactor.

The polymerization was allowed to proceed for the desired time and then the solution was drained from the bottom of the reactor and quenched with isopropanol. A hindered phenolic antioxidant (Irganox® 1010, available from Ciba-Geigy Corporation) was added and the polymer was air dried overnight. The residual solvent was removed in a vacuum oven overnight.

Polymer melt indices (I2) were measured according to ASTM D-1238, Procedure A, Condition E, using a micro-melt indexer, Model CS-127MF, available from Custom Scientific Instruments, Inc. Densities are determined by buoyancy measurements of compression molded samples in methyl ethyl ketone.

Experimental Determination of Lewis acidity

The Lewis acidity of phenylbis(perfluorophenyl)borane was determined using NMR analysis substantially according to the technique of R. Childs, et al., <u>Can. J. Chem.</u>, 1982, 60, 802-808. All manipulations were carried out either using standard Schlenk and/or high vacuum techniques or in an inert glovebox (Vacuum Atmospheres, Inc.) under recirculating, dry nitrogen with oxygen levels in the glove box being continuously monitored with an oxygen analyzer and maintained at < lppm. Crotonaldehyde purified by vacuum-transfer from calcium hydride and methylene chloride-d₂ (CD₂Cl₂) vacuum transferred from phosphorus pentoxide were used.

Proton NMR spectra were recorded either on a Varian VXR-300 NMR spectrometer or a Varian Gemini 300 NMR spectrometer. Chemical shifts were determined relative to tetramethylsilane through the residual CHDCl₂ (¹H, δ 5.32) in CD₂Cl₂. The NMR samples were prepared by adding an appropriate quantity of the Lewis acid compound to a solution of crotonaldehyde in CD₂Cl₂ at -20°C and allowing the sample to warm slowly to room temperature. The stoichiometry of the resulting solution was such that there was a 50 percent excess of the Lewis acid reagent with a total concentration of reactants being about 0.3M. The ¹H NMR spectrum was then recorded, and H-3 NMR shift of the Lewis acid adduct of crotonaldehyde and free crotonaldehyde determined.

Theoretical calculation of Lewis acidity

Structures and energies were calculated using one or more of the following standard techniques of lectronic structure theory.

1. AMI - Dewar's semi-empirical method based in approximate molecular orbital thirty. AMI has been par-

ametrized to fit selected experimental data. AMI calculati n techniques are well known t those skilled in th art, and are described in, for example, M. J. S. Dewar, E. G. Zoebisch, E. F. Healy, and J. J. P. Stewart, J. Am. Chem. Soc., 107, 3902 (1985); M. J. S. Dewar and Ev G. Zoebisch, J. M. I. Struct. (THEOCHEM) 180, 1 (1988); M. J. S. Dewar, C. Ji , and E. G. Zoebisch, Organ metallics, 7, 513 (1988); M. J. S. Dewar and C. Jie, Organometallics, 6, 1486 (1987); M. J. S. Dewar and K. M. Merz, Jr., Organometallics, 7, 522-4 (1988); M. J. S. Dewar and C. Jie, Organometallics, 8, 1547 (1989); M. J. S. Dewar and C. Jie, Organometallics, 8, 1544-7 (1989); and M. J. S. Dewar and A. J. Holder, Organometallics, 9, 508 (1990).

The AMI calculations are performed with the program MOPAC, version 5.00, available from the Quantum Chemistry Program Exchange (QCPE), Department of Chemistry, Indiana University, Bloomington, Indiana 47405. The program is further described in the MOPAC Manual, J. J. P. Stewart, Frank J. Seiler, Res. Lab., U.S. Air Force Academy, Colo. Spgs., CO. 80840

2. HF (restricted Hartree-Fock) the rigorous (ab initio, no adjustable parameters) method. The HF results were calculated using the GAUSSIAN® 90 program and the well-known 3-21g basis set. The 3-21g basis set is a valence double-zeta basis set. Gaussian 90, Revision J., available from Gaussian, Inc., Pittsburgh PA, 1990

The 3-21G basis set is well known in the art, and is described in, for example, W. J. Hehre, L. Radom, P. v. R. Schleyer, and J. A. Pople, *Abinitio Molecular Orbital Theory*, Wiley, New York, (1986); Pietro, W. J., et al., <u>J. Am. Chem. Soc.</u> 104, 5039-48 (1982); M. S. Gordon, et al., <u>J. Am. Chem. Soc.</u> 104, 2797-803 (1982); J. S. Binkley, et al., <u>J. Am. Chem. Soc.</u> 102, 939-47 (1980); K. D. Dobbs and W. J. Hehre, <u>J. Comput. Chem.</u> 8, 880-93 (1987); K. D. Dobbs and W. J. Hehre, <u>J. Comput. Chem.</u> 8, 861 (1987); K. D. Dobbs and W. J. Hehre, J. Comput. Chem. 7, 359 (1986).

Example 1

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A catalyst solution was prepared by combining 1 mL of a 0.005 M toluene solution of tris(penta-fluorophenyl)borane with 1 mL of a 0.005 M toluene solution of [(tert-butylamido)dimethyl(tetramethyl- η^5 -cyclopentadienyl)silane dibenzyl titanium [[(C_5Me_4)SiMe $_2$ N(t-Bu)]Tibz $_2$] (prepared by reaction of (tert-butylamido)dimethyl(tetramethyl)- η^5 -cyclopentadienyl)silane titanium dichloride and benzyl lithium. The dichloride in turn was prepared by reaction of lithium 1,2,3,4-tetramethylcyclopentadienide with (N-t-butylamino)(dimethyl)silane chloride, followed by conversion to the dilithium salt, reaction with TiCl $_3$ to form the closed ring structure (N-t-butylamido)dimethyl(tetramethyl- η^6 -cyclopentadienyl)titanium chloride, and oxidation of the metal center with methylene chloride to form (N-t-butylamido)dimethyl(tetramethyl- η^6 -cyclopentadienyl)titanium dichloride. The mixture was shaken for 10 seconds at 25°C, giving a catalyst solution noticeably darker than the starting, red-orange, titanium containing solution.

Polymerization

The catalyst solution was combined with a mixture comprising 1 L of Isopar E™, 0.2 L of 1-octene, and ethylene (31 atmospheres, 3.1 MPa) in a 2 L reactor. The reactants were previously deaerated and purified and the reactor contents were heated to 130°C. An immediate 7 °C rise in reactor temperature occurred. Ethylene was provided on demand at 3.1 MPa. After 10 minutes the reactor contents were removed and devolatilized giving 80.0 g of ethylene-octene copolymer having a micro-melt index of 0.104.

Example 2

A catalyst solution was prepared by combining 1 mL of a 0.005 M toluene solution of tris(pentafluorophenyl)borane with 1 mL of a 0.005 M toluene solution of (tert-butylamido)dimethyl(tetramethyl- η^5 -cyclopentadienyl)silane dibenzyl zirconium [(C₅Me₄)SiMe₂N(t-Bu)]Zrbz₂] (prepared according to the same technique as Example 1). The mixture was shaken for 10 seconds at 25°C, giving a catalyst solution slightly darker than the starting, light yellow, zirconium containing solution.

Polymerization

The catalyst solution (10 µmol in 2ml Isopar™ E) was combined with a mixture comprising 0.8 L of Isopar E™ and 0.4 liters of propylene in a 2 L reactor. The reactor contents were heated to 50°C. After 45 minutes the reactor contents were removed and devolatilized giving 30.1 g of polypropyl ne having a micro-melt index of 24.3 and a syndiotactic index (measured at a racemic triad) f 83.5.

Examples 3-37

Th metal compl x for Examples 3-37 was (tert-butylamid)dimethyl(tetramethyl-5-cyclopentadienyl)silan titanium dimethyl, $[(C_5Me_4)SiMe_2N(t-Bu)]TiMe_2$, and the Lewis acid was tris(perfluorophenyl)borane, $B(C_6F_5)_3$. The polymerization time for all examples was 10 minutes. The results for Examples 3-37 are shown in Table I.

10		Density, <u>R/mL</u>	ı	ł	ı	1	1	1	1	ı		ı	•		ı	ı	1	0.9220	0.9173	0.9115
15		MI (12)	37.9	27.54	0.04	5.5	10.6	21.23	5.23	. 1.6	1.99	3.54	5.34	0.2	4.34	0.04	•	3.54	1.04	1.35
20		Yield, g	15.1	18.7	6.3	25.8	30.2	8.6	14	7.1	9.05	17.1	16.7	26.2	20.4	28	1.1	22.4	45.5	77.3
25		Ti (uMoles)	10.00	00.9	6.00	6.00	4.00	10.00	6.00	10.00	4.00	00.9	00.9	4.00	00.9	4.00	10.00	1.25	1.25	1.25
30	Table I	Lewis acid (n Mole)	15	6	6	6	9	15	6	15	9	6	6	9	6	9	15	1.25	1.25	1.25
35	e i	l-octene (mL)		0	0	0	0	0	0	0	0	0	0	0	0	0	0	55	55	55
		MPa	45	45	45	45	45	45	45	45	45	45	45	45	45	45	45	28	28	28
45		C ₂ H ₄ MPa	ë.	3,45	'n	3.	e,	9	3.	e.	'n	3.	3.	ë	3.	ë	e,	3.	e e	m
50		H2 AkPa	300	340	0	170	300	170	170	20	170	170	170	20	170	0	0	340	170	170
		I(°C)	204	190	190	190	176	210	190	204	170	190	190	176	190	170	210	163	163	150
55		X	æ	4	5	9	7	6 0	6	10	=======================================	12	13	14	15	16	17	18	. 19	20

5		Density, <u>R/mL</u>	0.9189	0.9310	0.9077	0.9079	0.9190	0.9173	0.9125	0.9223	0.9116	0.9337	0.9154	0.9192	0.9206	9606.0	0.9214	0.9176	0.9266
10	٠	MI (12)	1.07	1.93	5.6	97.0	0.07	1.44	2.04	0.19	4.24	0.15	0.58	1.14	9.95	1.01	3.78	0.81	0.36
15		Yield, g	49.1	30.1	43.2	10.2	35.5	39.7	23.7	22.9	77	16.1	10.2	30.8	22.6	42	17.7	32.1	31.3
20		<u>Ti</u> (μMoles)	1.25	1.25	1.25	1.25	1.25	1.25	1.25	1.25	1.25	1.25	1.25	1.25	1.25	1.25	1.25	1.25	1.25
25	Table I (con't)	Lewis acid (u Mole)	1.25			ł	1.25	1.25	1.25	1.25	1.25	1.25	1.25	1.25	1.25	1.25	1:25	1.25	1:25
30	Table	l-octene l	55	28	100	55	28	55	82	28	82	10	55	55	82	82	55	55	28
35		C_2H_4MPa	3.28	3.28	3.28	3.28	3.28	3.28	3.28	3.28	3.28	3.28	3.28	3.28	3.28	3.28	3.28	3.28	3.28
40		$\underline{H}_2 \underline{\Delta k P a}$	170	280	170	170	69	170	69	=	280	170	0	170	280	69	170	170	280
45		T(°C)	163	170	163	163	155	163	170	170	155	163	163	163	170	155	175	163	155
50		EX	21	22	23	24	25	56	27	28	29	30	31	32	33	34	35	36	37

Examples 38 - 57

The catalyst and procedure of Examples 3-37 were used except that 1000 mL of Isopar® E was added to the react r followed by various amounts of propylen. The desired amount of hydrogen was then added and the solution was heated to the reaction temperature. The solution was then saturated with ethylene at 475 psig (3.38 MPa). Equimolar am unts of the metal complex. [(C₅Me₄)SiMe₂N(t-Bu)]TiMe₂ and the Lewis acid. B(C₆F₅)₃ were mix d in 2 mL of Isopar® E and this solution was transferred into the reactor to begin the poly-

merization. The reaction was allowed to proceed for 15 minutes with ethylene being supplied on demand at 475 psig (3.38 MPa). The propylene content of the polymer was determined by carbon NMR spectroscopy using the method of J.C. Randall, Rev. Macromo. Ch. m. Phys., 29(2&3), 201-317, (1989). The results ar shown in Tabl. II.

												,										
10		Density, (g/mL)	0.8520	0.8513	0.8522	0.8516	0.8514	0.8518	0.8516	0.8631	0.8544	0.8518	0.8540	0.8536	0.8501	0.8559	0.8491	0.8518	0.8533	0.8517	0.8497	0.8508
15		Propylene <u>Molž</u>	30.7	40.0	41.2	31.9	50.2	46.0	38.0	24.2	30.0	34.3	30.0	38.2	38.1	28.2	35.8	37.5	32.9	35.2	43.2	36.8
20		Melt Index (12)	3.551	0.671	0.184	0.063	6.984	0.956	0.777	0.372	1.098	0.261	2.122	4.093	1.203	0.000	0.796	0.674	0.830	0.733	1.697	0.503
25	II	Yield, R	70.9	109.0	98.1	11.7	99.5	92.8	88.7	79.1	94.1	96.8	75.6	91.6	96.5	72.8	79.2	82.3	97.6	82.5	84.1	83.1
30	Table	Ti (umol)	.500	.375	.325	.325	.750	.375	.375	.375	.375	.300	.500	.750	.500	.250	.325	.325	.750	.325	.500	.325
35		H2 AkPa	069	550	140	0	550	340	340	340	550	340	550	340	340	140	340	340	140	340	140	340
40		Temp (°C)	95	550	98	95	104	95	95	95	98	80	104	110	95	98	95	95	104	95	104	95
45		$\frac{Propene}{(g)}$	100	98	130	100	130	100	100	20	7.0	100	7.0	100	100	70	100	100	70	100	130	100
50		SI X	38	39	40	41	42	43	77	45	97	47	48	65	20	51	52	53	54	55	56	57

Examples 59 - 77

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The procedure of Examples 3 - 37 was followed except that various amounts of styrene were added to the react r followed by Isopar® E t make 1000 mL of total liquid volum. Various amounts of propyline were then added. The solution was heated to thi reaction temperature. This solution is then saturated with ethylene at 480 psig (3.41 MPa). Equimolar amounts of thim metal complex, [(C_5Me_4)SiM $_2N(t-Bu)$]TiMe $_2$, and the Lewis acid, B(C_6F_5)3 were mixed in 2 mL of Isopar® E and this solution was transferred into thin reactor to begin the

polymerization. The reaction was allowed to proceed for 10 minutes with ethylene being supplied on d mand at 480 psig (3.41 MPa). The results are shown in Table III. Where provided, propylene and styrene contents were determined by carbon 13 NMR spectroscopy.

5		Melt Index (12)	0.078	0.819	7.494	37.200	2.557	29.000	0.893	0.214	2.607	7.415	2.784	966.0	0.982	0.566	99.100	1.221	0.920	0.125	0.048	0.210	percent
		Yield, g	57.3	34.3	31.4	9.04	51.2	38.2	9.04	73.4	71.2	42.2	53.7	40.2	9.95	44.8	27.3	51.6	45.1	53.8	64.9	79.1	mole p
15		_																					7.7
	III	Ti (umol)	1.25	1.25	3.75	2.50	3.75	3.75	1.25	1.25	1.25	2.50	1.50	1.25	1.25	j.50	3.75	1.25	1.25	1.25	0.75	0.75	styrene
25	Table	Temp (°C)	82	100	118	. 118	100	118	100	82	100	118	100	100	100	001	130	100	100	. 82	70	82	percent,
		Styrene (mL)	121	225	329	121	400	329	225	329	20	121	225	225	225	225	225	225	225	329	225	121	mole
30																							30
35		Propylene (g)	20	100	70	130	100	130	100	130	100	70	150	100	100	20	100	100	100	70	100	130	Propylene
		X	28	29	09	61	62 *	63	64	65	99	19	89	69	70	7.1	12	73	74	75	76	11	* Pr

40 Examples 78-100

The procedure of Examples 3-37 was followed except that various amounts of styrene were added to the reactor followed by Isopar® E to make 1000 mL of total liquid volume. Various amounts of hydrogen were then added. The solution was heated to the reaction temperature. The solution was then saturated with ethylene at 475 psig (3.38 MPa). Equimolar amounts of the metal complex, [(C₅Me₄)SiMe₂N(t-Bu)]TiMe₂ and the Lewis acid, B(C₆F₅)₃ were mixed in 2 mL of Isopar® E and this solution was transferred into the reactor to begin the polymerization. Two additional charges of complex and Lewis acid were added at 5-10 minute intervals. The total amount of catalyst added (based on titanium) is shown in Table IV. The reaction was allowed to proceed for 10 minutes after the final catalyst addition with ethylene being supplied on demand at 475 psig (3.38 MPa) throughout the run. The results are shown in Table IV.

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5		Styrene Content (mole %)	8.0	0.5	1.5	1.7	1.3	1.4	1.5	1	1.9	1.2	•	1.0	1.6	•	1	8.0	1.4	•	•	•	•	9.0	1
10		Melt Index (12)	0.209	0.436	2.117	1.884	1.471	1.352	2.447	0.739	2.055	7.111	0.335	4.061	9.417	3.494	1.144	0.710	1.134	0.270	1.185	0.465	1.126	0.489	0.115
15 20	-	Yield, g	24.0	44.8	37.5	59.8	52.5	52.0	34.4	65.1	46.2	31.1	46.3	49.2	39.8	55.1	53.1	46.4	52.6	45.5	52.2	50.2	51.3	63.0	52.5
. 25	Table IV	Catalyst (µmol)	5.0	5.0	7.5	7.5	7.5	7.5	7.5	5.8	7.5	7.5	3.0	7.5	7.5	7.5	7.5	7.5	7.5	5.0	7.5	7.5	7.5	7.5	5.0
30		Hydrogen (kPa)	0	76	76	110	69	:	28	110	69	=	Ξ	=		140	69	28	69	28	69	0	69	11	28
35		Temp (°C)	145	145	145	136	145	145	154	136	145	160	130	154	154	145	145	154	145	136	145	145	145	145	136
40		Styrene (mL)	125	75	225	234	175	175	234	116	275	175	175	116	234	175	175	116	175	234	175	175	175	75	116
		KX	78	79	80	8.	83	83	84	82	86	8	88	83	8	16	92	93	76	95	96	97	86	66	100

5 Example 101

The polymerization procedure of Examples 3-37 was followed using 1.25 micromoles of (tert-butylamido)dimethyl(tetramethyl- η 5-cyclopentadienyl)silane titanium dibenzyl [(C_5Me_4)SiMe₂N(t-Bu)]Ti bz₂ and 1.25 micromoles of the Lewis acid, B(C_6F_5)₃ mixed in 2 mL of Isopar[®] E. The reaction temperature was 160°C. 10 g of propylene and 100 Δ psi (0.7 MPa) hydrogen were added. The ethylene pressure was 460 psig (3.3 MPa). The polymerization time was 15 minutes. 22.9 g of an ethylene / propylene copolymer were isolated.

Example 102

The polymerizati in procedur of Exampl 101 was follow d using 1.00 micromoles of {(C₅Me₄)SiMe₂N(t-Bu)]TiMe₂ and 1.00 micromoles of the Lewis acid B(C₆F₅)₃ mixed in 2 mL of Isopar® E. The reaction temperature was 90°C. 1000 mL if Isopar® E and 200 mL of 1-octen were charged into the react in and no hydrogen was added. The other pressure was 500 psig (3.55 MPa). The polymerization time was 15 minutes. 85.9 g of

an thylene/ octene copolym r of approximately 0.87 g/mL density and m It index (I2) of 0.3 was isolated.

Example 103

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Polymerization using (tert-butylamido)dimethyl(tetrahydrofluoren)silan titanium dimethyl metal derivativ

Preparation of tetrahydrofluorene

15g (90.2 mmol) of fluorene was dissolved in 200ml of a tetrahydrofuran (THF)/ ethylene diamine solution (1:1 ratio). The solution was cooled in a ice bath and with stirring 3.13g of lithium (451.2 mmol) was added in small portions. After all the lithium had been added the solution was stirred for 2 h with consequent dissolution of the lithium metal. The resulting solution was then poured into a HCl/ice mixture. The solution was then extracted with diethyl ether. The organic washings were combined, washed with water and dried over MgSO₄. The solution was filtered and the solvent removed on a rotary evaporator. The crude material was then purified by dissolving in hexane and passing through a silica gel column to give 11.4g (75 percent yield) of product after solvent removal.

Preparation of lithium tetrahydrofluorenide

10 g of tetrahydofluorene (59 mmol) was dissolved in 75 ml of pentane. To this solution 21 ml of n-butyl lithium (n-BuLi) (2.65 M) was added dropwise over a 20 min period. The solution was stirred overnight with deposition of a white solid. The solid was collected by filtration, washed with pentane and dried under reduced pressure to give 7.14g (70 percent yield) of product.

Preparation of (N-t-butylamino)(dimethyl)(tetrahydrofluorenyl)silane

5.78 g (34.9 mmol) of CISiMe₂NHCMe₃ (prepared according to the method described in J. Prakt. Chem, 24(3-4), 226-30 (1964)) was added to 100 ml of THF. To this solution 6.15 g (34.9 mmol) of lithium tetrahydrofluorenide was added. The solution was then brought to reflux for 10 min and the solution cooled to room temperature. Gas chromatographic (GC) analysis showed the reaction to be complete. The solvent was then removed under reduced pressure, the residue extracted with pentane, filtered and the solvent again removed under reduced pressure to give 9.80 g of product (94 percent yield).

Preparation of dilithium (N-t-butylamido)(dimethyl)(tetrahydrofluorenyl)silane

9.80g (32.8 mmol) of (N-t-butylamino)(dimethyl)(tetrahydrofluorenyl)silane was dissolved in 100ml of diethylether. To this solution 26.6 ml (70.6 mmol) of nBuLi (2.65 M) was added dropwise. Upon complete addition of the nBuLi the solution was stirred for 2 h whereupon the solvent was removed under reduced pressure to give an oily orange residue which solidified upon trituration with pentane to give 11.86 g (98 percent yield) of a yellow solid which was identified by nuclear magnetic resonance spectroscopy (NMR) as an etherate adduct (3/4 Et_2O per molecule) of dillthium (N-t-butylamino)(dimethyl)(tetrahydrofluorenyl)silane.

Preparation of [(N-t-butylamido)(dimethyl)(tetrahydrofluorenyl)silane]titanium dichloride ([(tetrahydrofluorenyl)SiMe₂N(t-Bu)]TiCl₂)

6 g (16.1 mmol) of TiCl₃(THF)₃ was dissolved in 75 ml of THF. To this solution 5.92 g (16.1 mmol) of dilithium (N-t-butylamido)(dimethyl)(tetrahydrofluorenyl)silane(3/4 Et₂O) was added as a solid with stirring. The solution was then stirred for 45 min, after this time period PbCl₂ (2.25g, 8.1 mmol) was added and the solution stirred for 45 minutes. The THF was then removed under reduced pressure. The residue was then extracted with toluene, the solution filtered and the toluene removed under reduced pressure. The residue was then triturated with pentane and the solution cooled to -20°C for 3 hours. The red precipitate was collected via filtration and washed with cold pentane and dried under vacuum to yield 5.00 g (75 percent yield) of product.

Preparation of [(N-t-butylamido)(dimethyl)(tetrahydrofluorenyl)silane]titanium dimethyl ([(tetrahydrofluorenyl) SiM ₂N(t-Bu)]Ti(CH₃)₂)

5.0g of [(N-t-butylamido)(dimethyl)(tetrahydrofluorenyl)silane]titanium dichloride (12 mmol) was suspended in 100 ml of Et₂O. T this suspensi in 8.40 ml of methyl magnesium iodid (MeMgl) (3.0 M in diethyl ether.

 $\rm Et_2O$) was added dropwise with stirring over a 20 min period. Upon completion of the addition of the MeMgl the solution was stirred for 40 minutes. After this time period the $\rm Et_2O$ was removed under reduced pressure and the residue extracted with pentane, the solution filtered and the filtrat evaporated to dryness under reduced pressure to giv 3.48 g (77 percent yield) of product.

Polymerization

The polymerization procedure of Example 102 was followed using 2.00 micromoles of [(tetrahydrofluorenyl)SiMe $_2$ N(t-Bu)]TiMe $_2$ and 2.00 micromoles of the Lewis acid, B(C $_6$ F $_5$) $_3$ mixed in 2 mL of Isopar $_6$ E. The reaction temperature was 130°C. 808 g of Isopar $_6$ E and 100 g of 1-octene are charged into the reactor. No hydrogen was added. The ethylene pressure was 500 psig (3.55 MPa). The polymerization time was 15 minutes. 41.1 g of an ethylene/ octene copolymer were isolated.

Example 104.

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The procedure of Example 103 was followed except that 2.50 micromoles of [(tetrahydrofluorenyl)SiMe $_2$ N(t-Bu)]TiMe $_2$ and 2.50 micromoles of the Lewis acid B(C $_8$ F $_5$) $_3$, were mixed in 2 mL of Isopar $_8$ E were used to form the catalyst. The reaction temperature was 150°C. 829 g of Isopar $_8$ E and 29 g of 1-octene are charged into the reactor and no hydrogen was added. The ethylene pressure was 500 psig (3.55 MPa). The polymerization time was 15 minutes. 11.4 g of an ethylene/ octene copolymer were isolated.

Example 105.

Polymerization using (tert-butylamido)dimethyl(1,3-dimethyl-5,6,7,8-tetrahydroindenyl)silane titanium dimethyl metal derivative

Preparation of 4,5,6,7-tetrahydro-1-methyl-indan-3-one

Cyclohexene (27.3g, 0.33 mol), crotonic acid (28.7g, 0.33mol) and polyphosphoric acid (300ml) were mechanically stirred under a nitrogen atmosphere at 60°C for 30 minutes. The slurry was poured into water and the aqueous solution extracted with diethyl ether. The diethyl ether extract was washed successively with a 10 percent NaHCO₃ solution and a saturated NaCl solution. The organic extract was then dried over anhydrous MgSO4. The solution was then filtered and the solvent removed under reduced pressure. The crude product was then purified via vacuum distillation (bp 87-92°C at 5 torr, 0.7 kPa) to give 32.6g (66 percent) of purified material.

Preparation of 7,9 dimethylbicyclo-[4.3.0]-nona-l(6),7-diene

Methyl lithium (1.5M, 96ml) was added dropwise to a solution of 4,5,6,7-tetrahydro-1-methyl-indan-3-one (17.7g, 0.118 mol) in 50ml of diethyl ether under an argon atmosphere whereupon the reaction mixture was refluxed for 18 hours. The mixture after this time period was hydrolyzed and the reaction mixture extracted with diethyl ether. The ether extracts were dried over anhydrous MgSO₄ and filtered. To the ether solution, 0.5ml of 6M HCl was added and the solution stirred for one hour. After this time period the ether solution was washed with water, dried over anhydrous MgSO₄, filtered and concentrated. Distillation at reduced pressure afforded 8.0g (45 percent) of product.

Preparation of lithium 1,3-dimethyl-5,6,7,8-tetrahydroindenide

7,9 dimethylblcyclo-[4.3.0]-nona-l(6),7-diene (5.0g, 33.5 mmol) was added to 100ml of pentane. To this solution n-BuLi in pentane (2.7M, 13ml) was added dropwise and the mixture stirred for 12 hours. The resulting white precipitate was collected via filtration, washed with pentane and dried under reduced pressure to give 5.02g (97 percent) of product.

Preparation of (N-t-butylamino)(dimethyl)(1,3-dimethyl-5,6,7,8-tetrahydroindenyl)silane

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0.77g of CISiMe₂NHCMe₃ (4.67 mmol) was added to 50 ml of THF. To this solution 0.75g (4.67 mmol) of lithium 1,3-dimethyl-5,6,7,8-t trahydroindenide was added. The solution was thin brought to reflux for 10 min and the solution was thin cooled to room temperature. Gas-Chromatogram (GC) analysis shi wild thin reaction

to b complete. The solvent was then removed under reduced pressure, the residue extracted with pentane, filtered and the solvent again removed under reduced pressure to give 1.21g of product (94 percent yield).

Preparation of dilithium (N-t-butylamido)(dim thyl)(1,3-dimethyl-5,6,7,8-tetrahydroindenyl)silan

1.21g (4.36 mmol) of (N-t-butylamino)(dimethyl)(1,3-dimethyl-5,6,7,8-tetrahydroindenyl)silane were dissolved in 100ml of diethylether. To this solution 5.72ml (9.15 mmol) of nBuLi (1.6 M in pentane) was added dropwise. Upon complete addition of the nBuLi the solution was stirred for 2 hours whereupon the solvent was removed under reduced pressure to give an oily yellow residue which solidified upon trituration with pentane to give 1.00g (79 percent yield) of a tan solid product.

Preparation of [(N-t-butylamido)(dimethyl)(1,3-dimethyl-5,6,7,8-tetrahydroindenyl)silane] titanium dichloride ([(1,3-dimethyl-tetrahydroindenyl)SiMe₂N(t-Bu)]TiCl₂)

0.64g (1.72 mmol) of TiCl₃(THF)₃ was dissolved in 75ml of THF. To this solution 0.50g (1.72 mmol) of dilithium (N-t-butylamido)(dimethyl)(1,3-dimethyl-5,6,7,8-tetrahydroindenyl)silane was added as a solid with stirring. The solution was then stirred for 45 min, after this time period PbCl₂ (0.239g, 0.86 mmol) was added and the solution stirred for 45 minutes. The THF was then removed under reduced pressure. The residue was then extracted with toluene, the solution filtered and the toluene removed under reduced pressure. The residue was then triturated with pentane and the solution cooled to -20°C for 3 hours. The product was collected via filtration and washed with cold pentane and dried under vacuum to yield 0.32g (47 percent yield) of product.

Preparation of [(N-t-Butylamido)(dimethyl)(1,3-dimethyl-5,6,7,8-tetrahydroindeny l)silane] titanium dimethyl ([(1,3-dimethyl-tetrahydroindenyl)SiMe₂N(t-bu)]Ti(CH₃)₂)

0.32g of (N-t-butylamido)(dimethyl)(1,3-dimethyl-5,6,7,8-tetrahydroindenyl)silane titanium dichloride (0.81 mmol) was suspended in 40 ml of Et_2O . To this suspension 0.56 ml of MeMgI (3.0 M in diethyl ether) was added dropwise with stirring over a 20 minute period. Upon completion of the addition of the MeMgI, the solution was stirred for 40 minutes. After this time period the Et_2O was removed under reduced pressure and the residue extracted with pentane, the solution filtered and the filtrate evaporated to dryness under reduced pressure to give 0.21g (73 percent yield) of product.

Polymerization

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The procedure of Example 103 was followed except that 0.50 micromoles of [(1,3-dimethyltetrahydroindenyl)SiMe₂N(t-Bu)]TiMe₂ and 0.50 micromoles of the Lewis acid B(C_6F_5)₃, mixed in 2 mL of Isopar® E were used to form the catalyst/ cocatalyst mixture. The reaction temperature was 120°C. 797 g of Isopar® E and 61 g of 1-octene are charged into the reactor and 20 Δ psi (0.14 Δ MPa) of hydrogen was added. The ethylene pressure was 500 psig (3.55 MPa). The polymerization time was 10 minutes. 29.2 g of an ethylene/ octene copolymer were isolated. The micro-melt index (I2), was 0.975.

Example 108.

The procedure of Example 105 was followed except that 0.10 micromoles of [(1,3-dimethyltetrahydroindenyl)SiMe₂N(t-Bu)]TiMe₂ and 0.10 micromoles of the Lewis acid, B(C_eF₆)₃ mixed in 2 mL of Isopar[®] E were used to form the catalyst mixture. The reaction temperature was 90°C. 715 g of Isopar[®] E and 143 g of 1-octene were charged into the reactor and 10 Δpsi (0.07 ΔMPa) of hydrogen was added. The ethylene pressure was 500 psig (3.45 MPa). The polymerization time was 10 minutes. 64.5 g of an ethylene/ octene copolymer were isolated. The melt index (I2) was 0.346.

Example 107.

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The procedure of Example 106 was followed except that 0.025 micromoles of [(C₅Me₄)SiMe₂N(t-Bu)]TiMe₂ and 0.025 micromoles of the Lewis acid, B(C₆F₅)₃ mixed in 2 mL of Isopar® E were used to form the catalyst. The reaction t imperatur was 50°C. 679 g of Is par® E and 179 g of 1-octene were charged into the reactor and 20 Δpsi (0.14 ΔMPa) of hydrogen was added. The ethylene pressure was 500 psig (3.55 MPa). The polymerization time was 60 minutes. 40.7 g of an thylene/ octen copolymer were isolated. The melt index, I2, was 0.166.

Example 108.

The procedur of Example 107 was followed except that 2.00 micromoles of [(tetrahydrofluorenyl)SiM $_2$ N(t-Bu)]Tibz $_2$ (prepar d by reacting [(N-t-butylamid)(dim thyl)t trahydrofluorenyl)silane] titanium dichloride with benzyl magnesium chloride) and 2.00 micromoles of the th Lewis acid, B(C_8F_5) $_3$ mixed in 2 mL of Isopar 8 E were used to form the catalyst. The reaction temperature was 150°C. 822 g of Isopar 8 E and 36 g of 1-octene were charged into the reactor and 10 Δ psi (0.07 Δ MPa) of hydrogen was added. The ethylene pressure was 500 psig (3.55 MPa). The polymerization time was 15 minutes. 20.1 g of an ethylene/ octene copolymer were isolated. The melt index, I2, was 0.327.

Example 109.

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The procedure of Example 108 was followed except that 2.00 micromoles of [(tetrahydrofluorenyl)SiMe₂N(t-Bu)]Tibz₂ and 2.00 micromoles of the Lewis acid, B(C₆F₆)₃ mixed in 2 mL of Isopar[®] E were used to form the catalyst The reaction temperature was 150°C. 822 g of Isopar[®] E and 36 g of 1-octene are charged into the reactor and 10 Δpsi (0.07 ΔMPa) of hydrogen was added. The ethylene pressure was 500 psig (3.55 MPa). The polymerization time was 15 minutes. 20.1 g of an ethylene/ octene copolymer were isolated. The melt index, 12, was 0.327.

20 Example 110.

Polymerization using (n-tertbutylamido)dimethyl(η^5 -tertbutylcyclopentadienyl)silane titanium dimethyl metal derivative

25 Lithium t-butylcyclopentadienide

To a solution of 4.18 g (39.4 mmol) 6,6-dimethylfulvene in 80 mL diethylether at 0°C were added 22.9 mL of a 1.72 M (39.4 mmol) methyl lithium solution in ether. The resulting slurry was stirred several days, then filtered, washed with pentane and dried under vacuum.

(N-t-butylamino)(dimethyl)(t-butylcyclopentadienyl)silane

To a solution of 3.35 g (20.2 mmol) (N-t-butylamino)(chloro)dimethylsilane in 75 mL THF was added 3.58 g (17.7 mmol) lithium t-butylcyclopentadienide etherate. The reaction mixture was stirred several hours. The solvent was removed, the residue was extracted with pentane and filtered. The pentane was removed in vacuo to give the product as a pale yellow oil. The yield was 2.87 g, 64.6 percent.

Dilithium[(N-t-butylamido)(dimethyl)(t-butylcyclopentadienyl)silane

To a solution of 2.87 g (11.4 mmol) (N-t-butylamino)(dimethyl)(t-butylcyclopentadienyl)silane in 70 mL ether was added 15.8 mL of 1.48 M (23.4 mmol) butyl lithium in hexane. The resulting clear solution was stirred overnight. The solvent was removed under reduced pressure. Yield was 107 percent of impure product.

[(N-t-butylamido)(dimethyl)(η-t-butylcyclopentadienyl)silane]titanium dichloride ([(t-butyl-C₅H₃)SiMe₂N(t-Bu)] TiCl₂)

In a flask were combined 0.60 g (2.27 mmol) dilithium[(N-t-butylamido)(dimethyl)(t-butylcyclopentadie-nyl)silane and 0.84 g (2.27 mmol) TiCl₃(THF)₃ as solids. To this was added 40 mL THF. The resulting dark violet solution was stirred for 10 minutes, then 0.35 g (1.25 mmol) PbCl₂ was added. The reaction mixture was stirred for less than one hour, the deep orange-brown reaction mixture was filtered, and the solvent removed under reduced pressure. The residue was extracted with pentane and the solution filtered and concentrated until solid material began to form. The slurry was cooled overnight in a freezer, the yellow product was collected on a frit, washed with pentane and dried under reduced pressure. The yield was 0.58 g, 69.6 percent.

55 [(N-t-butylamido)(dimethyl)(η-t-butylcyclopentadienyl)silane]titanium dimethyl ([(t-Butyl-C₅H₃)SiMe₂N(t-Bu)]Ti(CH₃)₂)

A 0.80 mL solution of 2.78 M (2.22 mmol) methyl magnesium iodide in 15 mL ether was slowly added over

20 minutes to 0.41 g (1.11 mmol) [(N-t-butylamid)(dimethyl)(η-t-butylcyclopentadienyl)silane]titanium dichloride in 15 mL ether. The solution was stirred 20 minutes, then the solvent was remov d. The residue was extracted with pentan and the resulting solution was filtered, then concentrated to an il which crystallized on standing. Yield was 0.34 g, 94.6 percent.

Polymerization

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The general polymerization procedure of Example 109 was followed using 0.25 micromoles of [(t-butyl- C_5H_3)SiMe₂N(t-Bu)]TiMe₂ and 0.25 micromoles of the Lewis acid, B(C_6F_5)₃ mixed in 2 mL of Isopar® E were used to form the catalyst. The reaction temperature was 80°C. 1000 mL of Isopar® E, 100 g of propylene and 50 Δ psi (0.34 Δ MPa) of hydrogen were added. The ethylene pressure was 475 psig (3.38 MPa). The polymerization time was 10 minutes. 6.3 g of an ethylene/ propylene copolymer was isolated. The melt index, I2, was 1.291. The density was 0.8868 g/mL.

5 Example 111. Ethylene / norbornylene copolymer

The general polymerization procedure of Example 109 was followed using 1.25 micromoles of [(C_5Me_4)Si-Me₂N(t-Bu)]TiMe₂ and 1.87 micromoles of the Lewis acid B(C_6F_6)₃ mixed in 2 mL of Isopar® E were used to form the catalyst. The reaction temperature was 140°C. 808 g of Isopar® E, 19.5 g of norbornene and 25 Δ psi (0.17 Δ MPa) of hydrogen were added. The ethylene pressure was 500 psig (3.55 MPa). The polymerization time was 10 minutes. 41.3 g of a randon ethylene / norbornene copolymer was isolated. The melt index (I2) was 0.587. The polymer contained 2.38 weight percent norbornene as determined by ¹³C NMR.

Example 112. Ethylene / norbornene copolymer

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The procedure of Example 111 was followed using 1.25 micromoles of [(C₅Me₄)SiMe₂N(t-Bu)]TiMe₂ and 1.87 micromoles of the Lewis acid, B(C₆F₅)₃, mixed in 2 mL of Isopar® E to form the catalyst. The reaction temperature was 140°C. 758 g of Isopar® E, 39.0 g of norbornene and 25 Δpsi (0.17 ΔMPa) of hydrogen were added. The ethylene pressure was 500 psig (3.55 MPa). The polymerization time was 10 minutes. 38.1 g of a random ethylene / norbornene copolymer were isolated. The melt index (I2) was 1.52. The polymer contained 4.33 weight percent norbornene as determined by ¹³C NMR.

Example 113. Ethylene / norbomene copolymer

The procedure of Example 112 was followed using 2.00 micromoles of $[(C_5Me_4)SiMe_2N(t-Bu)]TiMe_2$ and 3.00 micromoles of the Lewis acid, $B(C_6F_5)_3$, mixed in 2 mL of Isopar® E to form the catalysts / cocatalyst mixture. The reaction temperature was 50°C. 1200 mL of a solution containing 334.6 g of norbornene in Isopar® E and 5 Δ psi (0.03 Δ MPa) of hydrogen were added. The ethylene pressure was 100 psig (0.79 MPa). The polymerization time was 30 minutes. 22.9 g of a random ethylene / norbornene copolymer were isolated. The melt index, 12, was 1.43. The polymer contained 73.78 weight percent norbornene as determined by 13 C NMR. The polymer Tg was 83.8°C.

Example 114.

Polymerization using phenyl bis(perfluorophenyl)borane Lewis acid

Preparation of B(C₆F₅)₂(C₆H₅)

A 250 mL flask was evacuated, cooled to -78°C and charged with 120 mL of mixed hexane solvent, vacuum transferred from sodium benzophenone ketyl. The flask was backfilled with argon, to 0.11 MPa pressure, and bromopentafluorobenzene (deoxygenated by sparging with nitrogen, 10.00 grams, 40.5 mmol) was added via a syringe. Agitation of the mixture was begun (magnetic stir bar), giving a clear, colorless solution, to which was added n-butyllithium (16.2 mL of 2.5M hexane solution, 40.5 mmol) via syringe. As the n-butyllithium solution was added, a clear colorless solid separated from the mixture. This slurry was allowed to stir at -78°C for 70 minutes and then dichloroph nylboron (3.22g, 20.3 mmol, 0.50 quivalents) was added from a syring. After stirring for an additional 30 minutes at -78°C, no change was noted and the mixture were allowed to warm to ambient temperature. As the mixture warmed, a thick whit precipitate form d. After stirring at 22°C for 15 minutes th flask was open d to vacuum and th volum of th mixtur was reduced t 50 mL. The mixture was

filtered, the solid extracted with thre 20 mL portions if mixed he wans solvent and the filtration was reduced to 20 mL under reduced pressure. Their sulting solution was cooled to -78°C, giving a very thick slurry of colorless, crystalline solid. The slurry was thinned by addition of 20 mL if hexan in the solid was collected by filtration and dried under reduced pressure. Yield was 4.86g, 57 percent.

Polymerization

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The polymerization procedure of Examples 3-37 was followed except that 850 mL of Isopar® E is added to the reactor followed by 20 g of propylene. $25 \, \Delta psi$ (0.17 ΔMPa) of hydrogen were then added and the solution heated to $130^{\circ}C$. The solution was then saturated with ethylene at $500 \, psig$ (3.55 MPa). 10 micromoles of the metal complex [(C_5Me_4)SiMe₂N(t-Bu)]TiMe₂ and 10 micromoles of the Lewis acid, B(C_6F_5)₂(C_6H_6), were mixed in 2 mL of Isopar® E and this solution was transferred into the reactor to begin the polymerization. The reaction was allowed to proceed for 15 minutes with ethylene being supplied on demand at $500 \, psig$ (3.55 MPa). 2.8 g of an ethylene/ propylene copolymer were obtained. The melt index, I_2 , was 7.52.

Example 115 Ethylene / ethylidenenorbornene copolymer

The procedure of Example 111 was followed using two consecutive additions of a catalyst solution prepared by mixing 5.0 micromoles of $[C_5Me_4)SiMe_2N(t-Bu)]TiMe_2$ and 5.0 micromoles of the Lewis acid, $B(C_6F_5)_3$, mixed in 2 mL of Isopar® E. The reaction temperature was 130°C. 1200 mL of a solution containing 50 mL of 5-ethylidene-2-norbomene in Isopar® E and 50 psi (0.34 Δ MPa) hydrogen were added. The ethylene pressure was 475 psig (3.38 MPa). The polymerization time was 20 minutes. 59.9 g of an ethylene / 5-ethylidene-2-norbomene copolymer was isolated. The melt index, I2, was 1.55. The polymer contains 9.06 weight percent 5-ethylidene-2-norbenene as determined by 13 C NMR.

Example 116

Various Lewis acids are tested for Lewis acidity for use in preparation of catalysts according to the present invention. Acidity values and the techniques employed in determining such values are contained in Table V.

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Table V

5	Run	Lewis Acid	Acidity (Kcal/mole)	Base
	1	phenylbis(perfluorophenyl)borane	0.01,2,3	CH3-or NH3
	2	tris(2,3,5,6-tetrafluorophenyl)borane	-2.1 ²	CH ₃ -
10	3	tris(3,4,5-trifluorophenyl)borane	-5.2 ¹	•
	4	tris(3,4,5-trifluorophenyl)aluminum	-11.22	17
	5	tris(1,2,2-trifluoroethenyl)borane	-12.3 ¹	•
15	6	tris(2,3,4,5-tetrafluorophenyl)borane	-15.2 ²	*
	7	tris(perfluorophenyl) borate	-17.5 ¹	*
	8	tris(perfluorophenyl)borane	-17.81.5	
20	9	1,3,5-cyclohexanetriol borate	-22.2 ⁱ	NH ₃
	10	1,1,1-trimethylolethane borate	-25.1 ¹	п
25	1. 2. 3.	HF/3-21g method AM1 method Lewis acidity according to Childes' to ppm. Relative acidity = 0.0 ppm	echnique, $\Delta\delta$,	is 0.77
	4. 5.	B(OC ₆ F ₅) ₃ Lewis acidity according to Childes' to ppm. Relative acidity = 0.33 ppm	echnique, $\Delta\delta$,	is 1.10

Claims

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 A process for preparing a catalyst having a limiting charge separated structure corresponding to the formula:

LMX+ XA-,

wherein:

L is a derivative of a substituted delocalized Π-bonded group imparting a constrained geometry to the metal active site and containing up to 50 nonhydrogen atoms;

M is a metal of Group 4 or the Lanthanide series of the Periodic Table of the Elements;

X independently each occurrence is hydride or a hydrocarbyl, silyl or germyl group having up to 20 carbon, silicon or germanium atoms; and

A⁻ is an anion of a Lewis acid, A, having relative Lewis acidity greater than or equal to that of phenylbis(perfluorophenyl)borane, said anion being compatible with the metal cation,

the steps of the process comprising contacting a derivative of a Group 4 or Lanthanide metal corresponding to the formula:

LMX₂,

wherein

L, M, and X are as previously defined, with the Lewis acid, A.

- 2. A process as claimed in Claim 1 wherein M is titanium or zirconium.
- 3. A process according to Claim 1 wherein LMX₂ corresponds to the formula:

$$Cp \xrightarrow{Z} Y$$
 $(X)_2$

wherein:

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M is titanium or zirconium;

Cp is a substituted cyclopentadienyl group or derivative thereof that is Π -bound to M and substituted at least by Z;

Z is a divalent moiety comprising oxygen, sulfur, boron, or a member of Group 14 of the Periodic Table of the Elements;

Y is a ligand group comprising nitrogen, phosphorus, oxygen or sulfur or optionally Z and Y together form a fused ring system; and

X is as previously defined in claim 1.

- 4. A process as claimed in Claim 3, wherein -Z-Y- is an amidosilane or amidoalkane group.
- 5. A process as claimed in Claim 4, wherein LMX2 corresponds to the formula:

wherein:

M is titanium or zirconium, bound to an n⁶-cyclopentadienyl group;

R' each occurrence is independently selected from hydrogen, silyl, alkyl, aryl and combinations thereof having up to 20 carbon or silicon atoms, and optionally two or more R' groups on the cyclopenta-dienyl group may form a fused ring system;

E is silicon or carbon;

X independently each occurrence is hydride, alkyl, aryl or halogen substituted aryl of up to 20 carbons; and

m is 1 or 2.

- A process as claimed in Claim 5, wherein -(ER'₂)_m-NR'- has up to 20 nonhydrogen atoms.
 - A process as claimed in Claim 6, wherein -(ER'₂)_m-NR'- is (tert-butyl)(dimethylsilyl) or (tert-butyl)-1-ethan-2-yl.
- 45 8. A process as claimed in any one of the preceding claims, wherein X is hydride or C₁-C₁₀ hydrocarbyl.
 - 9. A process as claimed in Claim 8, wherein X is methyl or benzyl.
- 10. A process as claimed in Claim 9, wherein R' each occurrence in the cyclopentadienyl group is hydrogen or C₁-C₄ alkyl and optionally two or more of said R' together with the cyclopentadienyl group form a tetrahydroindenyl or tetrahydrofluorenyl group, and R' on the nitrogen atom is tert-butyl.
 - 11. A process as claimed in any one of the preceding claims wherein the Lewis acid is a boron compound lacking in halogen moieties directly attached to the boron.
- 12. A process as claimed in any one of the priceding claims, wherein the Lewis acid is selected from tris(pentafluorophenyl)borane, tris (2,3,5,6-tetrafluorophenyl)borane, tris(2,3,4,5-tetrafluorophenyl)borane, tris(3,4,5-trifluorophenyl)borane, tris(1,2,2-trifluoroethenyl)borane, phenylbis(perfluorophenyl)borane.

ane, tris(3,4,5-triflu rophenyl)aluminum, tris(perflu rophenyl) borat , 1,3,5-cyclohexanetriol borate, and 1,1,1-trimethylolethane borate.

- 13. A process as claimed in Claim 12, wherein th Lewis acid is tris(pentafluorophenyl)borane.
- 14. A process as claimed in any one of the preceding claims, wherein LMX $_2$ and A are contacted in a solvent selected from C_5 - C_{10} alkanes or cycloalkanes or mixtures thereof at 25°C to 50°C.
- 15. A process as claimed in any one of the preceding claims, wherein LMX₂ and A are contacted in situ during an addition polymerization reaction.
- 16. A process as claimed in any one of the preceding claims, wherein LMX₂ and A are contacted at an equivalent ratio of 0.5:1 to 1:2.
- 17. A catalyst obtainable by a process as claimed in any one of the preceding claims.

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- 18. The use of a catalyst as claimed in Claim 17 as an addition polymerization catalyst.
- 19. An addition polymerization process for preparing a polymer by contacting one or more addition polymerization catalyst under addition polymerization conditions, characterized in that the catalyst is as claimed in Claim 17.

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EUROPEAN SEARCH REPORT

Application Number

EP 92 30 5730

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